# **REMARKS**

In the Final Office Action dated June 21, 2006 the previously made restriction requirement was withdrawn. Accordingly claims drawn to additives containing calcium and magnesium ions, as well as claims to additives containing other polyvalent ions continue to be presented for consideration.

### Reorganization of the claims for re-examination

For the purpose of simplifying the presently requested Continuing Examination, the applicant, acting through the undersigned attorney, has canceled the original claims and submitted formal "new" Claims 21–48. To further simplify the task of examination, Claims 21-34 are drawn to the "calcium chloride, magnesium chloride and calcium oxide" subject matter. Claims 35 – 48 are drawn to the subject matter of "polyvalent ions,," now recited as Fe<sup>++</sup>, Fe<sup>+++</sup>, Cu<sup>++</sup> and Ba<sup>++</sup>.

The presently submitted claims represent editorial changes made for the purpose of placing the claims in better form, and also amendments of substantive subject matter which is fully supported by the original disclosure and avoids all art cited during the prosecution of this application.

# The rejection over prior art has been overcome or is inapplicable to the present claims

To show the *prima facie* allowability of all of the currently presented claims the applicant, acting through the undersigned attorney, discusses all pertinent references cited for anticipation or obviousness during the prosecution of this application, and shows that the present claims obviate all rejections based on these references.

Claims were rejected in the Office Action dated September 21, 2005 over published United States patent application US 2004/0058054 A1 (*Elder et al.*). This reference discloses

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reduction of acrylamide formation in thermally processed foods by adding calcium lactate, calcium citrate or calcium maleate to the food before thermal processing. The application also elaborates on the theory that acrylamide formation normally occurs because of decomposition or reaction of the amino acid asparagine in the food. Thus, the adding of calcium lactate, calcium citrate or calcium maleate to the food minimizes decomposition of asparagines to acrylamide during a thermal process.

Calcium lactate is only sparingly soluble in water. According to the enclosed *Handbook* of Chemistry and Physics 51<sup>st</sup> Edition (1970 – 1971) pp. B-77, 78, calcium lactate's solubility is 3.1g in 100 cc water. According to the same source, calcium citrate is even less soluble (0.85g in 100 cc of water) and calcium maleate has a solubility of 2.89g in 100 cc of water. Given the fact that in doughs, baked goods and the like the water content is relatively modest, it is unlikely that calcium lactate, calcium citrate or calcium maleate contribute a significant amount of calcium ions (Ca<sup>++</sup>) to the result described in the Elder et al. reference.

Even more importantly, the lactates, citrates and maleates or organic salts, each have the respective structural formulas shown below.

CH-OH 
$$H_2$$
C—COO  $H_2$ C—COO  $H_3$   $H_4$ C—COO  $H_4$ C—CO

Because the *Elder et al.* reference teaches that the decomposition or reaction of asparagines results in acrylamide formation and that addition of these organic salts minimizes such decomposition or reaction, a person of ordinary skill in the art is likely to conclude that it is the

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"organic" portion of these molecules which react or interact with asparagines and minimize the formation of acrylamide.

The present Claims 21-34 do not encompass the use of calcium lactate, citrate or maleate. Therefore any rejection for anticipation by  $Elder\ et\ al$ . is clearly avoided. For the reasons explained above a person of ordinary skill would not find it obvious to use calcium chloride, magnesium chloride or calcium oxide to minimize the formation of acrylamide. For this reason the subject matter of Claims 21-34 is not obvious over the  $Elder\ et\ al$ . reference or over any other reference of record.

"To serve as an anticipation when the reference is silent about the asserted inherent characteristic, such gap in the reference may be filled with recourse to extrinsic evidence. Such evidence must make clear that the missing descriptive matter is necessarily present in the thing described in the reference, and that it would be so recognized by persons of ordinary skill.:"

Continental Can Company USA, Inc. v. Monsanto Co., 948 F.2d 1264, 20 USPQ2d 1746 (Fed. Cir. 1991).

United States Patent No. 6,436,458 (*Kuechle et al.*) was cited in the Office Action of September 21, 2005. Applicant respectfully submits that this reference actually teaches away from the present claims. First, the main concern of *Kuechle et al.* is to provide a scoopable dough. The reference is not at all concerned and does not describe elimination or minimization of acrylamide formation during the cooking, baking or frying process.

Even more importantly, the reference contains several passages indicating that the presence of iron ions and other ions are to be avoided in the scoopable dough. Column 2, lines 6-9 of the reference state that the dough can include "flour enrichment with all <u>reduced iron</u>, a <u>binder of metal ions</u>, an organic acid and the like." (underlining added). Column 4, lines 5-6

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state that the scoopable dough "can include flour enrichments having iron that is substantially all reduced" (underlining added).

Column 10, lines 13 through 21 state:

"A scoopable dough of the invention can also include a <u>binder of metal ions</u>. Some metal ions can help catalyze enzymatic reactions that can result in dough graying. Thus, a binder of metal ions can be effective against a scoopable dough of the invention losing its natural color. Examples of binders of metal ions include <u>metal chelators and organic acids</u>. A metal chelator can include, for example, ethylene diaminotetracetate (<u>EDTA</u>)." (underlining added).

The term "all reduced iron" necessarily refers to metallic iron and not to iron ions. As is well known in the art "a binder of metal ion" is a chemical that removes the metal ion from effectiveness as an ion, by chelating it. EDTA is a very well known chelator of many kinds of metal ions, including that of iron ions and aluminum ions. Therefore, even if some of the "all reduced iron" in the *Kuechle et al.* disclosure were still in the ionic stage (two valent ferrous instead of three valent ferric), such ion would be removed by chelation. For all these reasons, the *Kuechle et al.* reference teaches away from the subject matter of the present Claims 35-48 and neither anticipates these claims nor renders them obvious.

United States Patent No. 5,556,655 (Vadlamani et al.) was cited in the Final Office Action of March 7, 2006. This reference is not concerned with eliminating or reducing the formation of acrylamide in foods which are subjected to thermal processes, such as cooking, baking or frying. Rather, its purpose is to reduce darkening of the food product during storage. The majority of examples of this reference are directed to a gluten product. The salts or ions added to the raw product (before it may be heated such as baked) were zinc and aluminum. The use of zinc and/or aluminum ions is no longer recited in the presently amended claims. Because

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the reference is entirely silent about reducing acrylamide formation, a person of ordinary skill in the art would receive no hint or suggestion that the use of Fe<sup>++</sup>, Fe<sup>+++</sup>, Cu<sup>++</sup> and/or Ba<sup>++</sup> ions, as stated in Claims 35 – 48, will reduce formation of acrylamides.

United States Patent No. 5,882,712 (*Wu*), cited in the Office Action of June 21, 2006, in spite of its appearance at first glance, does not disclose the use or presence of Ca *ions* originating from CaCl<sub>2</sub>, CaO or from other source of calcium, or of magnesium ions originating from MgCl<sub>2</sub> or from other source of magnesium while the therein disclosed dough or like batter product is baked or otherwise subjected to heat. On the contrary, this reference teaches to one of ordinary skill in the art that calcium or magnesium is no longer present in an ionic form, nor as CaCl<sub>2</sub> or MgCl<sub>2</sub> after the leavening process has been completed and the dough or batter is baked. The reasons are as follows.

The Wu reference discloses the use of a leavening composition for leavening a dough or batter before the dough or batter is baked or otherwise subjected to heat. The leavening composition of the reference includes a "leavening acid" and a "basic compound" which when reacted with the acid generates carbon dioxide. Carbon dioxide ( $CO_2$ ) is a gas that forms bubbles in the dough and therefore causes it to rise. This is well known in the art. It is significant that the reference refers to  $CaCl_2$  or  $MgCl_2$  as the "acid." In the ensuing discussion  $CaCl_2$  will be used as an example although the discussion is equally applicable to  $MgCl_2$ .

As a person of skill in the art will readily understand that carbon dioxide (CO<sub>2</sub>) gas forms in the reaction between CaCl<sub>2</sub> and the carbonate containing "basic" compound (such as sodium bicarbonate, NaHCO<sub>3</sub>) according to the following chemical equation.

$$CaCl_2 + 2NaHCO_3 \longrightarrow CaCO_3 + 2NaCl + H_2O + CO_2 + CO_3 + CO_3$$

As is well known in the art CO<sub>2</sub> is the gas that causes leavening of the batter, and calcium carbonate (CaCO<sub>3</sub>) is water insoluble (a substance that actually forms many mountain chains on the Earth). Water insoluble calcium carbonate does not contribute calcium ions (Ca<sup>++</sup>) to the batter like water soluble calcium chloride (CaCl<sub>2</sub>) does. The same type of chemical equation and discussion is applicable to water soluble magnesium chloride (MgCl<sub>2</sub>) and water insoluble MgCO<sub>3</sub>.

Because CaCl<sub>2</sub> and MgCl<sub>2</sub> is initially added to the batter or dough to leaven together with a basic compound (such as NaHCO<sub>3</sub>) and because, as shown above, calcium or magnesium ions are no longer present in substantive amounts in the batter or dough after it is leavened and while it is baked, the added calcium or magnesium of this reference does not inherently reduce acrylamide formation. Nor would a person of ordinary skill in the art find it obvious to omit the "basic compound" (such as NaHCO<sub>3</sub>) from the leavening substance added to the dough for the purpose of reducing acrylamide formation while the dough is baked.

The presently submitted independent Claims 21, 32 and 33 clearly set forth that the additive contributes calcium or magnesium ions to the dough or batter to be present while the dough or batter is cooked under heat. Therefore, the subject matter of Claims 21 - 34 clearly avoids the Wu reference.

United States Patent No. 4,272,554 (Schroeder et al.) cited in the Final Office Action of June 21, 2006 discloses preparing blister-inhibited potato chips. The disclosed process involves the step of applying small amount of calcium (a calcium chloride or other calcium salt) to the

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surface of the chips by "spraying, dipping or the like." Present claims 21-34 are not drawn to the making of fried or baked potato chips. This disclosure does not suggest to one of ordinary skill in the art that formation of acrylamide is reduced when calcium ions are added to a dough or batter. Therefore, the present claims clearly avoid the Schroeder et al. reference.

In light of the foregoing, all claims are free of the prior art and are in prima facie allowable condition. Their early allowance is respectfully solicited.

In the event the Examiner is of the opinion that a telephone conference with the undersigned attorney would materially facilitate the final disposition of this case, she is respectfully requested to telephone the undersigned attorney at the below listed telephone number.

I hereby certify that this correspondence is being Very truly yours, deposited with the U.S. Postal Service as first class mail in an envelope addressed to: Commissioner for Patent, PO Box 1450, Alexandria, VA 22313-1450 on August 31, 2006.

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# HANDBOOK CHEMISTRY and PHYSICS

# SIST

EDITION 1970-1971

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# PHYSICAL CONSTANTS OF INORGANIC COMPOUNDS (Continued)

	Name	Synonyms and Formulae	Mol. wt.	Crystalline form, properties and index of refraction	Density or spec. gravity	Melting point, °C	Boiling point, °C	Solubility, in grams per 100 cc				
o.								Cold water	Hot water	Other solvents		
$\dashv$	Calcium					;						
29	iodide, hexahydrate .	Cal <sub>2</sub> .6H <sub>2</sub> O	401.98	yel, hex need		d 42	160	7579	1680**	s a, al, acet		
30	iron (III) aluminate.	Calcium (tetra-)	485.97	brn, rhomb, 1.98,	3.77	1418 0				1.17	14.7	
ا ۳	HOII (III) aldiminate.	alumino-ferite, nat.		2.05, 2.08 all				3			OF	
		celite.		for \		l.	0	· · ·				
_]		4CaO.Fe <sub>2</sub> O <sub>2</sub> .Al <sub>2</sub> O <sub>2</sub>	:							i de la	1 to	
- 1		Ca(C4H7O2)2.5H2O	304.35	col powd				20	al a			
31	200000,1000	Ca(CaHiOi)1.5HiO	308.30	wh need, effl		-3H <sub>2</sub> O, 100	]	3.10	7.9*	alsa; i al, eth		
2	10000	Ca(C11401)1.5110	456.73	wh need, effl		182-183		0.00415	0.055100	0.059 <sup>18</sup> , 1.72 <sup>78</sup> al		
3			598.97	wh amorph powd				li.	:	a al, eth		٠.
4		Ca(C18H11O2)2	184.41	col, trig, 1.6817, ·		d 730-760		0.0321			2	
5	magnesium carbonate	Nat. dolomite.	101.11	1.5026							**************************************	
		CaCO <sub>3</sub> .MgCO <sub>3</sub>	216.56	col, monocl,	3.275	1391	·	i, 1	i	i HCl	is .	
88	magnesium	Nat. diopside.	210.00	1.665, 1.672,	0.5.0			1	j.	. 2		
i	metasilicate	CaO.MgO.2SiO <sub>1</sub>	٠.	1.695	1	1		1	i.s.	The Die		
۱			000 70		3.150			]		<u>2</u> 4		
7	magnesium	Nat. mervinite.	328.72	col to pa grn, monocl, 1.708,	10.100	1		1	1		a care	
- 1	orthosilicate	3CaO.MgO.SiO2			1		1	1	1.	P	<b>E</b>	
١		L	000 00	1.711, 1.718			1	0.3210	0.451#7.5	i al		
18	dl-malate	CaC4H4Os.3HsO	226.20	col, rhomb, 1.545,			1		,	7	E .	
١			.000 -0	1.555, 1.575			1	0.8120	1.224#.	a al	2	
9		CaC4H4Os.2HrO	208.18	col							<b>4</b>	
ю	malate, dihydrogen	Ca (HC4H4O4)2-6H2O	414.33	rhomb, or wh cr	1	1		[	f- ' '	5117		
١				powd, 1.493,	ļ		I "	į.			<b>S</b>	
- 1				1.507, 1.545	1		<b>J</b>	2.89**	3.21		Mary Street or other	
11	maleate	CaC4H1O4.H1O	172.15	col rhomb, 1.495,		1	1	15.00		Sea Sea	[4]	
		•		1.575, 1.640	1		l	0 440	0.72100		-	-
2	malonate	CaC <sub>1</sub> H <sub>2</sub> O <sub>4</sub> .4H <sub>2</sub> O	214.19						33825	a NH <sub>2</sub> OH	3	
3	permanganate	Ca(MnO <sub>4</sub> )2.5H2O	368.03	purp er		a			25.65 <sup>70</sup>		26	
4	a-methylbutyrate	Calvium ethylmethyl-	242.34					22.27	20.00			
-		acetate. Ca(CsHsO2)2				ļ,			d .	saial, eth	<b>3</b>	
15	molybdate	Nat. pawellite. CaMoO4	200.01	col, tetr, 1.967,	4.38-4.53	ļ		. ] 1	۳		·	
				1.978				121.2 <sup>18</sup>	376100	14" al; a MeOH, 15	£	
16	nitrate	Ca(NO <sub>1</sub> ) <sub>2</sub>	164.09	col, cub, hyg	2.504 <sup>11</sup>	561	ļ	121.2-	3,0	liq NH, acet; 9/9		
				1 :						i eth	Total and	
			ŀ		·.			0000	660**	s al, acet SOL	B	
47	nitrate, tetrahydrate	Ca(NO <sub>1</sub> )4.4H <sub>2</sub> O	236.15	col, monocl, deliq		a 42.7,	d 132	2660 .	000-	B al, acet		
*'	mulate, ventariyeraye	33,000	l	1.465, 1.498,	β 1.82	β 39.7			1		<b>3</b>	
				1.504					1 .			
48	nitrate, trihydrate	Ca(NO <sub>3</sub> )2.3H2O	218.14	col, tricl		51.1			1	s dil a; d abs al 🚉		
49		CasN <sub>1</sub>	148.25	brn cr, hex		1195			89.6 <sup>81</sup>	al a al		
50	nitrite	Ca (NO1)2. H2O	150.11	col-yelsh, hex,	2.23*	- H <sub>2</sub> O, 100		. 45.94	89.0-	81 8 21		
~	murae:			deliq						201		
			204.15	col cr, tetr	1.674	-2H <sub>2</sub> O, 44		74.90	10643	sal		
51	nitrite, tetrahydrate.	Ca(NO2)2.4H2O	204.15			. ,			0.0350	als eth	200	٠,
52	oleate	Ca (C18 H27O2)2	603.01	wh wax-like cr		83-84		0.042		sa; i ac a TOIS		•
53	oxalate	CaC2O4	128.10	col, cub		<b>d</b> '		0.000671	0.00148	sa; i ac a		
54	oxalate, hydrate	CaCrOs. HrO	146.12	col		- H <sub>2</sub> O; 200			2 000	- 3+200	5	
55	oxide	Lime, calcia. CaO	56.08	col, cub, 1.838 :	3.25-3.38	2580	2850	0.13110 d	0.07∞ d	S B	- Lander	
~	OAIGE			wh, tetr, 1.895	2 92 25	d 275		. sl s		. 8 a		
56	oxide, per	CaO2	72.08	Wn, tetr, 1.095	. 2.824			1 .	1.	NITY college in the	1	Ε,
.,		CaO2.8H2O	216.20	wh, tetr, pearly .	. 1.70	-8H <sub>2</sub> O, 200	d 275 expl	al s	d .	s a, NH salts; its		
57	oxide, per- octahydrate									i al, eth		
58	palmitate	Ca (C16H11O2)2	550.93	wh or yelsh, wh				0.003#		V 81 B A1, 0.000	The same	-
~	batimeng			fatty powd	1	1.	1 199			eth	42. 1	192
EC.	1-phenol-4	Ca[CeHe(OH)SOs]2.H2O	404.43	wh to pinkish		. [4	ļ	· la		s al		Ł.
59	sulfonate(p-)	02.00		powd	1 .			1.	.l.		- ** 24	4
	phenoxide	Ca (OC Hs):	226.29	redsh powd				. al s	1	. sla al		
60 ·	hypophosphate	Ca:P:Os.2H:O	274.13	gel		. :	.]	·   i	<b>!</b> }·····	s HCl	1	
61		Ca(PO <sub>2</sub> ) <sub>2</sub>	198.02	col, 1.588, 1.595.	. 2.82	975		· i · · · · · · ·	í	i a		-:
32	metaphosphate	Nat. brushite.	172.09	wh, tricl, 1.5576,	2.3061	-H <sub>2</sub> O, 109	[	. 0.0316 <sup>24</sup>	0.075100	i al, a a		
33	orthophosphate, di-(sec)	CaHPO.2HrO		1.5457, 1.5392	1		1	1	1	and Series		-
		Ca(H2PO4)2.H2O	252.07	col, tricl, deliq,	2,22016	-H <sub>2</sub> O, 109	d 203	1.820	d ·	5 8		
64	orthophosphate,			1.5292, 1.5176,			1		1			•
	mono-(prim.)		ľ	1.4392		1	1	l	ŀ.	530		
	authoritaritate .	Nat. whitlockite.	310.18	wh amorph powd	1, 3.14	1670		. 0.002	d	i al; s a		
65	orthophosphate,	Cas(PO <sub>4</sub> ):		1.629, 1.626	1	1	1	1	]	antes.	1	
	tri-(tert.)	Ca <sub>2</sub> P <sub>2</sub> O <sub>7</sub>	254.10	col, biax, 1.585,	3.09	1230		iji 🐪	· [,	. 8 8		
66	pyrophosphate	Carrio	202.10	1.604	1		1					
		O- D-O- 5H-O	344.18	col, monocl, 1.539	2.25	1	.	. al s	·	. s a; i NH <sub>4</sub> Cl	-11	
67	pyrophosphate.	CaiPiO1.5HiO	344.18	1.545, 1.551	'			1, 11	1			
	pentahydrate	'		gray lumps	. 2.51	ca 1600	1	d ev PH	1	sa; i al, eth, bs		
68	phosphide	CasP2	182.19		I	la	1	15.42	12.5100	ial west		
69	hypophosphite	Ca(ll2PO2)2	170.06	wh-gray, monocl	1	1	.1	al s	d	s NH <sub>6</sub> Cl	W 15 ~ .	6
70	orthophosphite, di	2Ca HPO:.3H:O	294.17	and be as	5.71	la	1	. li	ā	. 8 8		ď
	orthoplumbate	CasPbO	351.35	red-br cr		Ĭ	. [	490	55.8100	ial Sil	A 3	× 1
71		Ca.(CaHaO2)2.H2O	201.24	col, monocl tabl	4	111111111111111111111111111111111111111	3	1619	1	i al		
72	propionate	Ca(C7H11O1)1.10H1O	602.56	rhomb leaf		. 50, -10Hx	Д					

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# PHYSICAL CONSTANTS OF INORGANIC COMPOUNDS (Continued)

	1 - 35	· · —	<del></del>			IN 15 OF INON			(		• •	
		. No	o. Name	Synonyms and	Mól.		Density or	Melting		Solubility, in grams per 100 cc		
solvents		· _		Formulae	wt	index of refraction	gravity	point, °C	point, °	C Cold	1	Other solvents
		c c c	Calcium butyrate	Ca(C4H7O2)2.3H2O CaC2	268.3			stab	2300	s d	al a	
; s HCl		, c8	g carbonate	Nat. aragonite. CaC	Oa. 100.0	9 col, rhomb, 1.530, 1.681,	2.930	25-447 tr to calcite 520	d 825	0.00153	25 0.001907	s a, NH <sub>4</sub> Cl
; s exp INOs		<b>c</b> 9	0 carbonate	Nat. calcite. CaCO.	100.0	1.685 col, rhomb or hex	2.71018	13391025	d 898.6	0.001425	0.001875	s a, NH <sub>4</sub> Cl
h, glycerol		c9	carbonate, hexa- hydrate	CaCO <sub>1.6</sub> H <sub>2</sub> O	208.18	1.460, 1.535,	1.7710					
юн		c9	2 chiorate	Ca(ClO <sub>3</sub> ) <sub>2</sub>	206.99	1.545 wh cr, hyg		340 ± 10 (-some O		s	a	s al, acet
		c9	chlorate, dihydrate	Ca(ClO <sub>3</sub> ) <sub>2</sub> .2H <sub>2</sub> O	243.01	wh-yelsh, rhomb, or monocl, deliq	2.711	- H:O,	"	177.74	v s	s al, acet
		c9	perchlorate	(CaClO <sub>4</sub> ) <sub>2</sub>	238.98	col cr	2.651	d 270		188.62	v s	166.2 al; 237.4 MeOH
et, NHa		. c9!			110.99	col, cub, deliq	2.154	772	>1600	74.520	1591∞	s al, acet, ace a
		. c96			561.33	col, monocl or hex, hex, 1.550, 1.535 col cr	1.892 <sup>11</sup> 0.835	− H <sub>2</sub> O, 105	−8H <sub>2</sub> O, 35		d	s a
I . NHOH		c98		CaCls.6HsO		col, trig, deliq,		29.92	-4H <sub>2</sub> O, 30 -6H <sub>2</sub> O,	97.7° 279°	326₩ 536™	50 <sup>so</sup> al s al
i, NH1OH I1OH INOi		c99	hydrate	CaCl <sub>2</sub> .H <sub>2</sub> O				260	200		249100	s al; i acet
ЭН		: _ c101	orthophosphate chlorite	Ca(ClO <sub>2</sub> ) <sub>2</sub>	174.98	1.631 wh, cub	3.14 2.71	1270		v sl s	d	i al
one H		c102		Ca(ClO)2	1	pl. 1.545, 1.69	2.35 2.10	d 100		s sl s solns	d	i al d a
NHs;	7 E.S.			,		1.585				with 5-6 % avail		u.2
; i bs		c104	hypochlorite, basic.	Bleaching powder, chlorinated lime. Ca(ClO) <sub>2</sub> .CaCl <sub>2</sub> .xCa(O	comp varies:	wh powd strong Cl odor		d		d evin Ci		d a
i HNO <sub>1</sub> ,		c105	hypochlorite, tri- hydrate chromate	Ca (CIO)2.3 H2O	197.03	tetr pl, 1.535, 1.63 2	1	-311₂O, 60				
4,,		c107	chromite	CaCrO4.2HrO. CaCr2O4. Ca(CeHrO2)2.3H2O	208.07	yel, monocl pr	.814 2			i		ss, al ia; sfus K2CO2
		c109 c110	citrate	Ca <sub>2</sub> (C <sub>6</sub> H <sub>6</sub> O <sub>7</sub> ) <sub>2</sub> 4H <sub>2</sub> O CaCN <sub>2</sub>	80.10	wh need		-4H <sub>2</sub> O, 120		0.854	0.962	0.0065 <sup>12</sup> al
		cl11 cl12	, k	Ca(CN) <sub>2</sub>	92.12 429.31	wh powd		l >350 -5H±O, 100	• • • • • • • • • • • • • • • • • • • •	d . s	d ,.	.,
y		c113 c114	ferrite, mono	Cas[Fe(CN)4]212H2O CaO.Fe2O2	760.42 215.77	red need, deliqdk redsh r, rhomb, 5, 2,58, 2,43 (Na)	.08	250		v s i	v s i	v sl s a
		ci 15		Ca:Fe(CN)ell or 12[[20]		yel tricl, 1.570, 1.582, 1.596	.68 . d					al.
h HNO: al; i eth		c117	fluoride	Nat. fluorite. CaF2	182.16 78.08	I	.66 <sup>10</sup>	360		sl s 0.0016 <sup>18</sup>	0.001726	al, HF, HCl NH4 salts; sl s a; i acet
		c118 c119	formate	CaSiFe.2H2O Ca(CHO2)2	218.19 130.12	col, tetrag 2.: col, rhomb, 1.510, 2.: 1.514, 1.578				sl s d 16.2°		HCl, HF; i al al
H4 salts;		c120 c121 c122	d-gluconate glycerophosphate	CaC4H2O4.3H2O Ca(C4H11O1)2.H2O CaC2H4(OH)2PO4	208.18 448.40 210.16	col, rhomb		H <sub>2</sub> O, 120		2.1120 3.315		slsal
		c123	hydride	СаН2	42.10	wh, rhomb cr 1.9	9 . 81 . d	6 (in H <sub>2</sub> ) coi 600	d	H <sub>2</sub> + Ca(OH) <sub>2</sub>	d	al a
		c125		CaN201.4H2O	172.15	-1.545 wh cr	334. d:	H <sub>2</sub> O, 580 d	d	0.1850		NH. salts, a; i al dil a
Oz; al s conc			iodata					1	- 1-			
O <sub>1</sub> ; als conc O <sub>4</sub> acet, a; als		c126	iodate	Nat. lautarite. Ca(IO <sub>2</sub> ) <sub>2</sub> Ca(IO <sub>2</sub> ) <sub>2</sub> ,6H <sub>2</sub> O				540				HNO: i al
Os; al s conc O4		c126	iodateiodate, hexahydrate	Ca(IO2)2.6H2O	497.98	col, rhomb	519 <sup>13</sup> d ;	35	o	.130	.22 <sup>100</sup> s	HNO; i al HNO; 26 <sup>20</sup> MeOH; s al, acet, a

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